Please amend the above-referenced application as follows:

In The Specification:

Please replace the Sequence Listing (1 page) filed on April 19, 2002 with the substitute Sequence Listing (1 page) filed herewith.

Please replace the paragraph beginning at page 6, line 5, with the following rewritten paragraph:

As used herein, "analyte" refers to any atom and/or molecule; including their complexes and fragment ions. In the case of biological molecules/macromolecules or "biopolymers", such analytes include but are not limited to: proteins, peptides, DNA, RNA, carbohydrates, steroids, and lipids. Note that most important biomolecules under investigation for their involvement in the structure or regulation of life processes are quite large (typically several thousand times larger than $\rm H_2O$).

Please replace the paragraph beginning at page 19, line 2, with the following rewritten paragraph:

FIGURE 1 is a representation of derived data which characterizes a disease specific marker having a particular sequence (SEQ ID NO:1) useful in evidencing and categorizing at least one particular disease state 7. Each patient listed in the data table shows the presence of the disease specific marker (SEQ ID NO:1) in their serum.

Please replace the paragraph beginning at page 19, line 6, with the following rewritten paragraph:

FIGURE 2 is the characteristic profile derived via SELDI/TOF MS of the disease specific marker of Figure 1. SEQ ID NO:1 is shown.

Please replace the paragraph beginning at page 22, line 19, with the following re-written paragraph:

Chelating Sepharose SEPHAROSE Mini Column

- 1. Dilute Sera in Sample/Running buffer;
- 2. Add Chelating Sepharose SEPHAROSE slurry to column and allow column to pack;
 - 3. Add UF water to the column to aid in packing;
- 4. Add Charging Buffer once water is at the level of the resin surface;

- 5. Add UF water to wash through non bound metal ions once charge buffer washes through;
- 6. Add running buffer to equilibrate column for sample loading;
 - 7. Add diluted serum sample;
 - 8. Add running buffer to wash unbound protein;
 - 9. Add elution buffer and collect elution fractions for analysis;
 - 10. Acidify each elution fraction.

Please replace the paragraph beginning at page 27, line 17, with the following re-written paragraph:

As a result of these procedures, the disease specific marker identified by the sequence SSKITHRIHWESASLLR (SEQ ID NO:1) was found. This marker is characterized as a C3f fragment from the complement system having a molecular weight of about 2056 daltons. The characteristic profile of the marker is set forth in Figure 2. As easily deduced from the data set forth in Figure 1, this marker is indicative of an individual suffering from congestive heart failure.

Please replace the paragraph beginning at page 36, line 2, with the following re-written paragraph:

The instant invention involves the use of a combination of preparatory steps in conjunction with mass spectroscopy and time-of-flight detection procedures to maximize the diversity of biopolymers which are verifiable within a particular sample. The cohort of biopolymers verified within such a sample is then viewed with reference to their ability to evidence at least one particular disease state; thereby enabling a diagnostician to gain the ability to characterize either the presence or absence of said at least one disease state relative to recognition of the presence and/or the absence of said the biopolymer.